§ 1002.2

TABLE 1—RECORD AND REPORTING REQUIREMENTS BY PRODUCT—Continued

Manufacturer							Dealer & Distributor
Products	Product reports § 1002.10	Supple- mental reports § 1002.11	Abbre- viated re- ports § 1002.12	Annual reports § 1002.13	Test records § 1002.30(a) ¹	Distribution records § 1002.30(b) ²	Distribution records §§ 1002.40 and 1002.41
Ultrasonic therapy (1050.10) Diagnostic ultrasound	Х	Х	х	Х	Х	Х	Х
Medical ultrasound other than therapy or diagnostic Nonmedical ultrasound	X	X	x				

¹However, authority to inspect all appropriate documents supporting the adequacy of a manufacturer's compliance testing program is retained.

[60 FR 48382, Sept. 19, 1995; 61 FR 13423, Mar. 27, 1996]

§1002.2 [Reserved]

§ 1002.3 Notification to user of performance and technical data.

As authorized by §5.90 of this chapter, the Director and Deputy Director of the Center for Devices and Radiological Health may require a manufacturer of a radiation emitting electronic product to provide to the ultimate purchaser, at the time of original purchase, such performance data and other technical data related to safety of the product as the Director or Deputy Director finds necessary.

[60 FR 48385, Sept. 19, 1995; 61 FR 13424, Mar. 27, 1996]

§ 1002.4 Confidentiality of information.

The Secretary or his representative shall not disclose any information reported to or otherwise obtained by him, pursuant to this part, which concerns or relates to a trade secret or other matter referred to in section 1905 of title 18 of the United States Code, except that such information may be disclosed to other officers or employees of the Department and of the other agencies concerned with carrying out the requirements of the Act. Nothing in this section shall authorize the withholding of information by the Secretary, or by any officers or employees under his control, from the duly authorized committees of the Congress.

§1002.7 Submission of data and reports.

All submissions such as reports, test data, product descriptions, and other information required by this part, or voluntarily submitted to the Director, Center for Devices and Radiological Health, shall be filed with the number of copies as prescribed by the Director, Center for Devices and Radiological Health, and shall be signed by the person making the submission. The submissions required by this part shall be addressed to the Center for Devices and Radiological Health, Electronic Product Reports, Office of Compliance (HFZ-307), 2098 Gaither Rd., Rockville, MD 20850.

- (a) In addition to the requirements of this part, all material submitted to the Director, Center for Devices and Radiological Health, shall be submitted pursuant to the provisions of part 20—Public Information, of this chapter.
- (b) Where guides or instructions have been issued by the Director for the submission of material required by this part, such as test data, product reports, abbreviated reports, supplemental reports, and annual reports, the material submitted shall conform to the applicable reporting guides or instructions. Where it is not feasible or where it would not be appropriate to conform to any portion of a prescribed reporting

ram is retained.

2The requirement includes §§ 1002.31 and 1002.42, if applicable.

3Report of Assembly (Form FDA 2579) is required for diagnostic x-ray components; see 21 CFR 1020.30(d)(1) through (d)(3).

4Systems records and reports are required if a manufacturer exercises the option and certifies the system as permitted in 21

Determined using the isoexposure rate limit curve (IRLC) under phase III test conditions (1020.10(c)(3)(iii)).

Annual report is for production status information only.

Determination of the applicable reporting category for a laser product shall be based on the worst-case hazard present within the laser product.